Heterogeneous and Dynamic Prevalence of Asymptomatic Influenza Virus Infections

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Influenza infection manifests in a wide spectrum of severity, including symptomless pathogen carriers. We conducted a systematic review and meta-analysis of 55 studies to elucidate the proportional representation of these asymptomatic infected persons. We observed extensive heterogeneity among these studies. The prevalence of asymptomatic carriage (total absence of symptoms) ranged from 5.2% to 35.5% and subclinical cases (illness that did not meet the criteria for acute respiratory or influenza-like illness) from 25.4% to 61.8%. Statistical analysis showed that the heterogeneity could not be explained by the type of influenza, the laboratory tests used to detect the virus, the year of the study, or the location of the study. Projections of infection spread and strategies for disease control require that we identify the proportional representation of these insidious spreaders early on in the emergence of new influenza subtypes or strains and track how this rate evolves over time and space.

Infection of the respiratory tract with an influenza virus results in symptoms ranging from mild nonfebrile illness to severe disease and complications, including pneumonia, shock, renal failure, encephalopathy, and multiorgan dysfunction (1,2). Influenza viruses infect 5%–15% of the global population annually (3), accounting for \approx 500,000 deaths (4) and 19 million disability-adjusted life years (5). The occurrence of asymptomatic influenza viruses infections has been recognized for some time (6), but determinations about their possible role in transmission are largely speculative (7,8). Clarifying the role of these infections in virus transmission requires a solid understanding of their rate of occurrence.

Interest in the contribution of asymptomatic infection to influenza virus transmission has risen in recent years

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after a series of outbreaks caused by newly emerging subtypes (9–12). Subclinical infection eludes symptomatic surveillance, and resulting illnesses thus manifest as sporadic disease. Social network analysis indicates that nearly one third of the attack rate for influenza A(H1N1)pdm09 virus in England was attributable to asymptomatic infection (13), a proportion mirrored by a recent review of volunteer challenge studies (14). Mathematical modeling studies designed to inform pandemic preparedness and vaccination thresholds and stockpiling strategies have typically had to resort to using these types of indirect metrics for parameterization (15–17). Current policy surrounding intervention planning for pandemic and interpandemic influenza is informed by estimates and simulations that arbitrarily assume a constant rate of asymptomatic infection in the range of 30%–50%.

However, mortality rates, clinical symptoms, and basic reproduction numbers (outbreak thresholds) vary greatly between influenza virus types, subtypes, and strains (18). Therefore, assigning an arbitrary value for asymptomatic infection rates that does not reflect this heterogeneity presents an important shortcoming in the current ability to accurately predict influenza outbreaks. Therefore, we conducted a systematic review and meta-analysis to determine the prevalence of asymptomatic influenza infection and to identify any factors associated with the heterogeneity reported across studies.

Methods

Search Strategy and Selection Criteria

A systematic review and meta-analysis was conducted in accordance with PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) guidelines (19). Literature searches were performed on the PubMed and Web of Science databases for the period from the inception of these databases to the beginning of 2015 to identify studies that reported laboratory-confirmed influenza infection (i.e., by culture, PCR, or serologic testing) and the proportion of symptomatic versus asymptomatic presentation. Search terms were chosen to ensure maximum coverage of possible literature and included the terms "influenza,"

"carrier," "carriage," "shedding," "asymptomatic," "influenza AND prophylaxis NOT vaccine" (filtered for randomized control trials), "influenza AND (travel OR migration OR immigra*) AND (screening OR test OR testing OR detection)," "subclinical," "serosurvey OR seroprevalence OR seroepidemiology." Other keywords and connectors were also used (online Technical Appendix 1, http://wwwnc.cdc.gov/EID/article/22/6/15-1080-Techapp1.pdf).

To be eligible for inclusion, studies needed to 1) be peer-reviewed and 2) report the prevalence of asymptomatic influenza virus infections in humans or present the appropriate data from which that prevalence could be calculated. Laboratory confirmation of influenza was a requirement, and it had to be possible to correlate these data to the number of symptomatic patients. We did not impose limitations in terms of study design, influenza virus type, or exposure type (community or experimental inoculation). According to current World Health Organization guidelines, laboratory confirmation consisted of 1) conventional PCR (referred to here as PCR) or real-time reverse transcription PCR (rRT-PCR); 2) virus antigen detection by immunofluorescence or enzyme immunoassay methods; 3) serologic detection of antibodies (hemagglutination inhibition); or 4) virus culture (20). Studies were excluded when the use of antiviral agents without a placebo group was reported. In cases in which a placebo group was used and an asymptomatic proportion could be determined, only this subset was used; otherwise, the study was excluded. Results were restricted to studies published in English; however, no restriction was placed on the publication date of studies that fit these criteria.

Study Selection and Data Extraction

Two authors (L.F-K. and M.C.) independently screened the publications for eligibility in a stepwise fashion. Search results were initially screened based on article titles and abstracts. Then, full-text analysis was performed to identify all studies which either reported asymptomatic prevalence or from which asymptomatic prevalence could be calculated. Any discrepancies that might have affected inclusion or exclusion of a study were resolved through discussion and consensus after independent evaluation by another author (L.Y.). The same 2 authors (L.F-K. and M.C.) assessed the risk for bias of the studies included by using a modified version of the tool developed by Hoy et al. (21) for prevalence studies (online Technical Appendix 2, http://wwwnc.cdc.gov/EID/article/22/6/15-1080-Techapp2.pdf).

The definitions of asymptomatic influenza infection varied considerably between studies. Definitions ranged from a total absence of symptoms to a lack of influenza-like illness (ILI) or acute respiratory illness (ARI). For the sake of clarity, we used the term "asymptomatic" when there was a total absence of symptoms and "subclinical" when the patient did not meet the authors' criteria for ILI

or ARI. Asymptomatic influenza prevalence was considered to be the proportion of all persons with laboratory-confirmed influenza who had no symptoms, whereas subclinical influenza prevalence was the proportion of persons with laboratory-confirmed influenza who failed to meet the study's definition of symptomatic infection. In addition to collecting data on asymptomatic and subclinical infection prevalence, we collected data on influenza virus type/subtype and study characteristics (e.g., study design, sample size, diagnostic test used to detect influenza virus infection, and the working definition of "symptomatic").

Statistical Analysis

We used prevalence of asymptomatic versus subclinical carriers among persons with laboratory-confirmed influenza as primary endpoints of interest. We pooled the prevalence estimates of asymptomatic and subclinical influenza across studies by using 2 meta-analytical models, the inverse variance heterogeneity model (22) and the random effects model.

We observed considerable heterogeneity across studies. This heterogeneity was unlikely to be attributable only to random or systematic errors, and actual clinical heterogeneity was deemed to exist. Therefore, we created subgroups by influenza virus type/subtype with the aim of generating more homogeneous groups within which we could anticipate that the differences indeed reflected variability caused by random or systematic error rather than actual clinical heterogeneity. In addition, we built a linear model to examine the variance explained by the influenza virus type/subtype, laboratory test used to detect the virus, year of the study, and geographic location of the study to gain insight into the considerable heterogeneity observed in the prevalence of asymptomatic and subclinical infections. We conducted the meta-analyses by using MetaXL version 2.0 (EpiGear Int Ptv Ltd, Brisbane, OLD, Australia), which also included the inverse variance heterogeneity method, and the generalized linear model by using Stata version 12 (StataCorp LP, College Station, TX, USA). All tests were 2-tailed, and a p value < 0.05 was deemed statistically significant.

Results

Yield of Search Strategy

A total of 13,219 records were identified from literature searches of the 2 databases. This number was reduced to 9,900 after removal of publications that were either duplicates or not original research papers (e.g., review papers). An additional 3,663 papers were removed based on the title and 5,652 papers more based on the abstract. The full texts of the remaining 585 studies were examined, and 55 articles met the inclusion criteria and were included in the final analysis (Figure; online Technical Appendix 1).

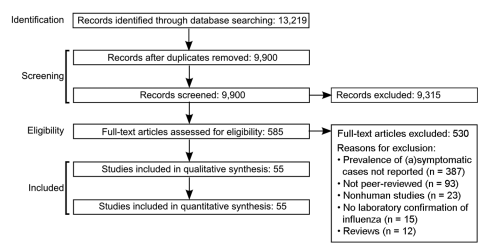


Figure. PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analysis) flowchart of literature search for systematic review and meta-analysis of asymptomatic and subclinical influenza infection prevalence.

Characteristics of the Studies Included

The 55 articles provided 59 data points because 4 papers reported the prevalence of asymptomatic and subclinical carriers for influenza A and B viruses separately. Overall, 19 studies (22 data points) defined asymptomatic infection as cases in persons lacking symptoms, and 44 studies (46 data points) reported subclinical influenza virus infections.

Infection was confirmed by serologic testing, rRT-PCR, or viral culture; 28 studies reported use of serologic testing alone to confirm infection, 18 used rRT-PCR alone, and the remaining 9 used a combination of methods (5 serologic testing and rRT-PCR, 3 serologic testing and culture, and 1 rRT-PCR and culture). Among the 55 studies, influenza A virus (predominantly H1N1) was the most common type of infection; 5 studies reported influenza B virus infections, and 1 study reported influenza C infections (online Technical Appendix 1 Table 2). Most studies reported on pandemic influenza virus types; 32 of these studies related to the 2009 pandemic influenza A/Mexico/4108/2009 strain. The risk for bias was moderate in 32% of the studies and low in the remaining 68%; no study was found to have a high risk for bias.

Quantitative Synthesis

The overall pooled prevalence for asymptomatic carriers was 19.1% (95% CI 5.2%–35.5%) for any type of influenza,

21.0% (95% CI 4.2%–41.0%) for influenza A, and 22.7% (95% CI 7.7%–39.8%) for influenza A(H1N1) (Table 1; online Technical Appendix 2 Figure 1). For subclinical carriers, the overall pooled prevalence was 43.4% (95% CI 25.4%–61.8%) for any type of influenza, 42.8% (95% CI 22.3%–63.9%) for influenza A, and 39.8% (95% CI 16.4%–64.5%) for influenza A(H1N1) (Table 1; online Technical Appendix 2 Figure 2). However, extensive heterogeneity was immediately evident for reported asymptomatic prevalence ($\tau^2 = 0.31$) and subclinical prevalence ($\tau^2 = 0.45$) that could not be explained by the influenza type/subtype alone. Similar results were obtained with the random effects model (online Technical Appendix 2 Figures 3, 4).

Investigation of Heterogeneity

The considerable heterogeneity observed within asymptomatic and subclinical influenza prevalence could not be explained by the type/subtype of influenza, the laboratory tests used to detect the virus, the location where the study was conducted, or the year of the study. The multivariate regression models could only explain 16.8% and 14.8% of the observed variance for the asymptomatic and subclinical prevalence, respectively. Influenza type/subtype as an independent predictor was found to account for almost the entire variance (16%) found for the prevalence of asymptomatic carriers (Table 2).

Table 1. Heterogeneity within asymptomatic and subclinical influenza infection cases, by virus type/subtype, as determined through a systematic review and meta-analysis of 55 studies

Type/subtype	Prevalence (95% CI)	Cochran's Q	p value (Cochran's Q)	l ² ,* %
Asymptomatic				
All types of influenza	19.1 (5.2–35.5)	752.40	<0.001	97
Influenza A	21.0 (4.2–41.0)	692.94	<0.001	98
Influenza A(H1N1)	22.7 (7.7–39.8)	561.14	<0.001	97
Subclinical	· · ·			
All types of influenza	43.4 (25.4–61.8)	1768.24	<0.001	97
Influenza A	42.8 (22.3–63.9)	1689.78	<0.001	98
Influenza A(H1N1)	39.8 (16.4–64.5)	1388.54	<0.001	98

^{*}The I² statistic describes the percentage of variation across studies that is attributable to heterogeneity rather than chance.

Table 2. Variance attributable to predictors in univariate and multivariate regression models for asymptomatic and subclinical influenza infection prevalence, by study characteristics, as determined through a systematic review and meta-analysis of 55 studies

Model/characteristic	Asymptomatic	Subclinical
Univariate model		
Influenza type/subtype	0.1599	0.0345
Laboratory test used to detect influenza	0.0043	0.0546
Hemisphere where study was conducted	0.0001	0.0159
Continent where study was conducted	0.0045	0.0213
Decade when study was conducted	*	0.0064
Multivariate model	0.1676†	0.1478‡

^{*}Variance not reported because all the studies were from the same decade.

Publication Bias

For both asymptomatic and subclinical carrier prevalence, the funnel plots showed no indication of publication bias. This result was confirmed by Doi plots (data not shown).

Discussion

Studies of laboratory-confirmed influenza typically do not include details of the symptomatic versus asymptomatic rate of infection. Of the few that do include this information, ambiguity exists between definitions of asymptomatic versus subclinical infections. This has perpetuated the ubiquitous issue of absent denominators in documented influenza rates and has caused substantial aberrations in initial reports of newly emerging subtypes and strains (23). We propose that the term "asymptomatic" be used exclusively to describe the complete absence of symptoms associated with influenza virus infection in patients with laboratory-confirmed cases. Given that reporting of this rate in the clinical literature would require little to no additional effort for most study designs. we also propose that the asymptomatic rate of laboratory test-positive persons be declared explicitly by public health bodies and researchers.

We found no evidence to support a fixed asymptomatic rate (or even an informative range) between or even within influenza virus subtypes. For example, the prevalence of asymptomatic influenza A(H1N1) virus ranged from 0% to 65%, resulting in an overall failure to explain the extreme heterogeneity in this reported rate. Some alternative explanations for the extreme heterogeneity are plausible, one being that generally applicable biologic mechanisms underlie the asymptomatic rates of influenza virus infection and these have been missed (e.g., details of patient vaccination or infection history were not routinely described in the clinical studies and data on sex and age of patients were excluded). Alternatively, influenza viruses conferring asymptomatic infection mutate so rapidly that a meaningful single per-influenza type rate simply does not exist. Employing sensitive diagnostic testing and standardized reporting of the asymptomatic rate of influenza virus infection would elucidate any underlying mechanisms or demonstrate any temporal changes in this rate.

This lack of a convenient asymptomatic rate poses a considerable obstacle to public health planning. Disease surveillance and control strategy is contingent on reliable estimates for the asymptomatic rate and the contribution that asymptomatic persons have on transmissibility. For example, a low asymptomatic rate improves the utility of passive (i.e., symptom-based) surveillance, whereas a higher asymptomatic rate might prompt presumptive travel restrictions to curb the spread of newly emerging subtypes and strains, especially if a high mortality rate is evident early in the outbreak. Future analyses correlating asymptomatic rates with mortality rates are also required; although one could easily speculate that influenza subtypes and strains eliciting high asymptomatic rates probably incur correspondingly low mortality rates, no evidence supporting this assumption currently exists.

Our study clearly demonstrates the inappropriateness of a one-size-fits-all approach to mitigating the spread of human influenza viruses. As new subtypes and strains emerge, actively surveying infection status of local populations and tracking any changes in asymptomatic rates of infection should increasingly become a global health priority, possibly necessitating the provision of international resources and the deployment of dedicated rapid-response teams who are guided by standardized protocols.

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Mr. Furuya-Kanamori is an infectious disease epidemiologist. He is enrolled in a PhD program at the Australian National University where he uses modern quantitative methods to better understand risk factors associated with different infectious diseases.

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[†]Model adjusted for influenza type/subtype, laboratory test, and location (continent) of the study.

[‡]Model adjusted for influenza type/subtype, laboratory test, location (continent) of the study, and decade when the study was conducted.

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etymologia

Etymology is concerned with the origin of words, how they've evolved over time, and changed in

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featured monthly in

http://wwwnc.cdc.gov/eid/articles/etymologia INFECTIOUS DISEASES

EMERGING

Heterogeneous and Dynamic Prevalence of Asymptomatic Influenza Virus Infections

Technical Appendix 1

Search Strategies

First Search

Influenza AND ((Asymptomatic OR Carrier OR carriage OR shedding OR symptomatic OR Subclinical OR serosurvey OR seroprevalence OR seroepidemiology) OR ((travel OR migration OR immigra*) AND (screening OR test OR testing OR detection)) OR (("Cohort Studies" [Mesh] OR "Case-Control Studies" [Mesh]) AND "Influenza A virus" [Mesh]))

Second Search

("influenza, human" [MeSH Terms] OR ("influenza" [All Fields] AND "human" [All Fields]) OR "human influenza" [All Fields] OR "influenza" [All Fields]) AND ("prevention and control" [Subheading] OR ("prevention" [All Fields] AND "control" [All Fields]) OR "prevention and control" [All Fields] OR "prophylaxis" [All Fields]) NOT ("vaccines" [MeSH Terms] OR "vaccines" [All Fields] OR "vaccine" [All Fields])

Manually filtered for randomized controlled trials.

Subclinical Influenza type Seasonal / Definition of Definition of Asymptomatic prevalence, Authors, year Location of the study or subtype Pandemic Exposure type Diagnosis test asymptomatic subclinical prevalence, % Aho M, et al., 2010 A (H1N1) 40.7 Pandemic Military garrison No symptoms Finland HI ≥10 of URT infection

Technical Appendix Table 1. Characteristics of the 55 studies included in systematic review and meta-analysis of asymptomatic and subclinical influenza infection prevalence

						infection			
Belderok SM, et al., 2013	Netherlands	A, and B	Seasonal	Travel to tropical and subtropical countries	HI ≥40 and ≥4-fold increase above pre-travel titer		No ILI		90.83
Bone A, et al., 2012	France	A (H1N1)	Pandemic	Community	HI ≥40	_	No ILI		29.52
Buescher, et al., 1969	Thailand and Panama	A (H3N1)	Pandemic	Military garrison	HI ≥32		No ILI		77.5
Carey DE, et al., 1958	USA	A (H2N2)	Pandemic	Parish	HI ≥10	-	No 'flu'	-	24.77
Ceyhan M, et al., 2010	Turkey	A (H5N1)		Community, poultry exposure and healthcare workers	HI ≥21		No symptoms of avian influenza infection		81.25
Clover RD, et al., 1986*	USA	A (H1N1)	Seasonal	Community	Positive culture or HI ≥4-fold increase		No ILI		60
Cui F, et al., 2011	China	A (H1N1)	Pandemic	Train	rRT-PCR	_	No ARI	_	13.64
Dotan A, et al., 2014	Israel	A (H1N1)	Pandemic	Hospital	rRT-PCR	_	No URI	_	30.77
Du Ry van Beest Holle M, et al., 2005	Netherlands	A (H7N7)		Poultry	HI ≥10		No ILI		93.94
Foy HM, et al., 1987	USA	В	Seasonal	Community	HI ≥10	_	No ILI	_	32.43
Gray GC, et al., 2014	Cambodia	A (H1N1), A (H3N2), and B	Seasonal and pandemic	Community	HI ≥4-fold increase		No ILI	_	64.44
Guinard A, et al., 2009	France	A (H1N1)	Pandemic	School	rRT-PCR		No ILI		53.33
Hayden FG, et al., 1999*	USA	A (H1N1)	Seasonal	Experimental inoculation	Positive culture and/or HI ≥4-fold increase		No URT illness	_	46.15
Hayward AC, et al., 2014	UK	A (H1N1), A (H3N2), and B	Seasonal and pandemic	Community	rRT-PCR		No ILI	_	46.22
Hsieh YH, et al., 2014	Taiwan	A (H1N1)	Seasonal	Community and school	HI ≥4-fold increase	No symptoms	No ILI	45.15	33.33
Hudson L, et al., 2013	New Zealand	A (H1N1)	Pandemic	Healthcare workers	HI ≥40	None influenza symptoms	_	25.44	_
Ison MG, et al., 2012*	Belgium, Estonia, France, Germany, Hungary, Israel, Italy, Lithuania, Spain, UK, USA	A (H1N1), A (H3N2), and B		Transplant recipients	Positive culture and/or HI ≥4-fold increase	No symptoms		25	
Jackson ML, et al., 2011	USA	A (H1N1)	Pandemic	School	HI ≥20 and ≥4-fold increase	No symptoms	No ILI	25	81.25

Authors, year	Location of the study	Influenza type or subtype	Seasonal / Pandemic	Exposure type	Diagnosis test	Definition of asymptomatic	Definition of subclinical	Asymptomatic prevalence, %	Subclinical prevalence, %
Jaeger JL, et al., 2011	USA	A (H1N1)	Pandemic	Hospital	HI ≥20		No ARI or ILI		66.66
Johnson S, et al., 2011	UK	A (H1N1)	Pandemic	Boarding school	HI ≥8		No ILI	—	68.35
Khakpour M, et al., 1969	Iran	A (H3N2)	Pandemic	Prisoners	HI	—	No ILI	—	23.53
Khaokham CB, et al., 2013	USA	A (H1N1)	Pandemic	Navy vessel	rRT-PCR or HI ≥4- fold increase	No symptoms	No ILI	52.11	88.03
Khuntirat B, et al., 2014	Thailand	A (H1N1)	Pandemic	Community	rRT-PCR and HI ≥4-fold increase	-	No ILI	_	83.33
Kumar S, et al., 2010	USA	A (H1N1)	Pandemic	Community	rRT-PCR	No symptoms	No ILI	10	32
Kumar S, et al., 2011	USA	A (H1N1)	Pandemic	Healthcare workers	HI ≥40	No symptoms	No ILI	35	30
Kuster SP, et al., 2013	Canada	A (H1N1)	Pandemic	Community and healthcare workers	HI ≥40		No ARI	_	13.04
Lau LLH, et al., 2010	Hong Kong	A (H1N1), A (H3N2), and B	Seasonal	Community	rRT-PCR	No symptoms	_	25.42	
Levy JW, et al., 2013	Thailand	A (H1N1), A (H3N2), and B	Seasonal and pandemic	Community	rRT-PCR	No symptoms	_	2.54	_
Li T, et al., 2011	China	A (H1N1)	Pandemic	Boarding school	rRT-PCR and HI ≥40	No symptoms	_	30.89	_
Mikulska M, et al., 2013	Italy	A (H1N1), A (H3N2), and B	Seasonal	Allogeneic haematopoietic stem cell recipients	rRT-PCR	No symptoms	No ILI	10	45
Neatherlin J, et al., 2013	USA	A (H1N1)	Pandemic	Airplane	MN ≥40 and HI ≥20	—	No ARI/ILI	—	75
Oker-Blom N, et al., 1970*	Finland	A (H3N2)	Pandemic	Community	HI ≥4-fold increase	_	No respiratory illness	_	18
Pang X, et al., 2011	China	A (H1N1)	Pandemic	Community	rRT-PCR	_	No ILI		4.62
Papenburg J, et al., 2010	Canada	A (H1N1)	Pandemic	Community	Microneutralizatio n ≥40 or ≥4-fold increase	No symptoms	_	9.43	_
Pascalis H, et al., 2012	Reunion Island	A (H1N1)	Pandemic	Community	rRT-PCR	No symptoms	No ILI	1.61	30.65
Pasco JA, et al., 2012	Australia	A (H1N1)	Pandemic	Community	HI ≥40	_	No ILI	_	75.97
Paton NI, et al., 2011*	Singapore	A (H1N1), A (H3N2), and B	Seasonal	Community	HI ≥4-fold increase		No clinical influenza (ILI)		51.72
Priest PC, et al., 2013	New Zealand	A, and B	Seasonal	Airport	rRT-PCR	No symptoms	— (. <u></u>	6.67	_
Qi W, et al., 2014	China	A (H10N8)	Pandemic	Poultry exposure	HI ≥20	_	No influenza symptoms		100
Redlberger-Fritz M, et al., 2014	Austria	A (H1N1)	Pandemic	Attended hospital	rRT-PCR	_	No respiratory symptoms	_	60.72

Authors, year	Location of the study	Influenza type or subtype	Seasonal / Pandemic	Exposure type	Diagnosis test	Definition of asymptomatic	Definition of subclinical	Asymptomatic prevalence, %	Subclinical prevalence %
Robinson JL, et al., 2007	Canada	A (H3N2)	Tanacimo	Community	HI ≥32	<u>—</u>	No ILI	—	77.78
Salez N, et al., 2014	France, Reunion Island, UK	С		Community	HI, ELISA and rRT-PCR	-	No ILI	_	50
Shafir SC, et al., 2011	USA	A (H1N1)		University campus	HI ≥40	_	No ILI	-	54.43
Shankar AG, et al., 2014	UK	A (H1N1)	Pandemic	Airplane	rRT-PCR		No ILI		0
Smit PM, et al., 2012	Netherlands	A (H1N1)	Seasonal	Healthcare workers	rRT-PCR	_	No ILI	-	0
Sridhar S, et al., 2014	UK	A (H1N1)	Pandemic	Community	HI ≥32	_	No ILI	-	84.15
Suess T, et al., 2012	Germany	A (H1N1), A (H3N2), and B	Seasonal and pandemic	Community	rRT-PCR	No symptoms	No ILI	4.76	17.99
Thai PQ, et al., 2014	Vietnam	A (H1N1)	Pandemic	Community	rRT-PCR	No symptoms		45.45	
Toyokawa T, et al., 2011	Japan	A (H1N1)	Pandemic	Healthcare workers	HI ≥40		No fever		92.86
Vilella A, et al., 2012	Dominican Republic	A (H1N1)	Pandemic	Community	rRT-PCR	No symptoms		5.13	
Wang TE, et al., 2010	Taiwan	A (H1N1), A (H3N2)	Seasonal	School	HI ≥4-fold increase	No symptoms	_	62.5	_
Woods CW, et al., 2013	USA and UK	A (H1N1), A (H3N2)	Seasonal	Experimental inoculation	Positive culture or rRT-PCR		Jackson score <6	_	56.1
Yan L, et al., 2012	China	A (H1N1)	Pandemic	School	rRT-PCR or HI ≥40	-	No ARI	-	64.49
Zaman M, et al., 2011	Pakistan	A (H5N1)	Pandemic	Hospital	rRT-PCR	No symptoms		25	

^{*}Only control or placebo group included; ARI, acute respiratory illness; HI, hemagglutination inhibition; ILI, influenza-like illness; LRT, lower respiratory tract; rRT-PCR, real-time reverse transcription PCR; URT, upper respiratory tract.

Technical Appendix 2 Table. Included studies (N = 55), by influenza type/subtype

Туре	Subtype	No. of studies	References
Α	H10N8	1	1
	H7N7	1	2
	H5N1	2	3,4
	H3N2	11	5–15
	H2N2	1	16
	H1N1	38	5,8,9,12–15,17–47
В		5	5,8,12,48,49
С		1	50
Mixed		6	<i>4</i> 9, <i>51</i> – <i>55</i>

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Heterogeneous and Dynamic Prevalence of Asymptomatic Influenza Virus Infections

Technical Appendix 2

Risk for Bias Assessment

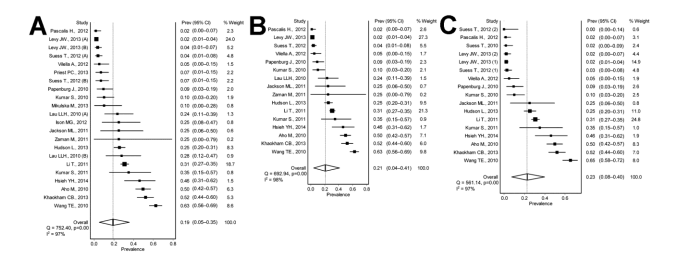
Risk for bias of the studies was assessed by using a modified version of the tool developed by Hoy et al. for prevalence studies. The modified tool assessed the external and internal validity of the studies by 9 criteria: 1) the targeted population was a close representation of the national population; 2) the sampling frame was a true or close representation of the general population; 3) random selection was used to select the study population; 4) the likelihood of nonresponse bias was minimal; 5) the case definition of influenza infection was based on laboratory tests; 6) the data collected were reliable; 7) the method used to collect the data was the same for all subjects; 8) the numerator and denominator of the prevalence were based on all the study participants; and 9) the data were largely recorded directly from the participants.

Statistical Analysis

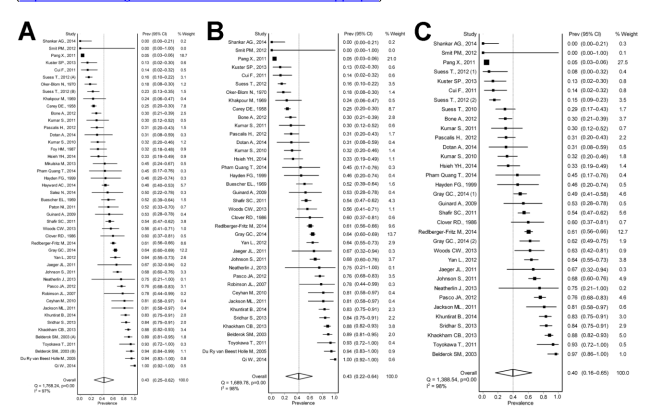
All prevalence estimates were pooled after double arcsine square root transformation and back-transformed for reporting (1). We did not use the standard random effects (RE) model (2) to pool because the latter is known to underestimate the statistical error and exacerbate publication bias (3–5). Therefore, the prevalence rates of asymptomatic and subclinical influenza across studies within the subgroups were pooled using the inverse variance heterogeneity (IVhet) model (6). This method uses a quasi-likelihood based variance structure without distributional assumptions and thus has coverage probabilities for the CI well within the 95% nominal level and has been documented to have better performance (lower mean squared error) when compared to the RE method (7). The results from the RE model have nevertheless been reported (Technical Appendix 2 Figures 3 and 4) for comparison purposes. Publication bias was assessed through Egger's linear regression and visual inspection of funnel and Doi plots (8).

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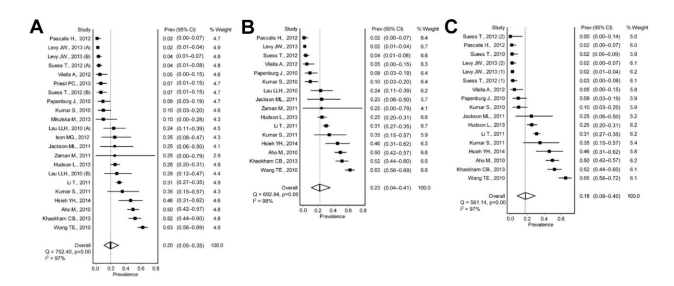


Technical Appendix 2 **Figure 1.** Forest plots for asymptomatic prevalence of A) influenza overall, B) influenza A, and C) influenza A(H1N1) virus infections generated by using the inverse variance heterogeneity model for 55 studies included in systematic review and meta-analysis of asymptomatic and subclinical influenza infection prevalence. Details on these studies are provided in Technical Appendix 1 (http://wwwnc.cdc.gov/EID/article/22/6/15-1080-Techapp1.pdf).

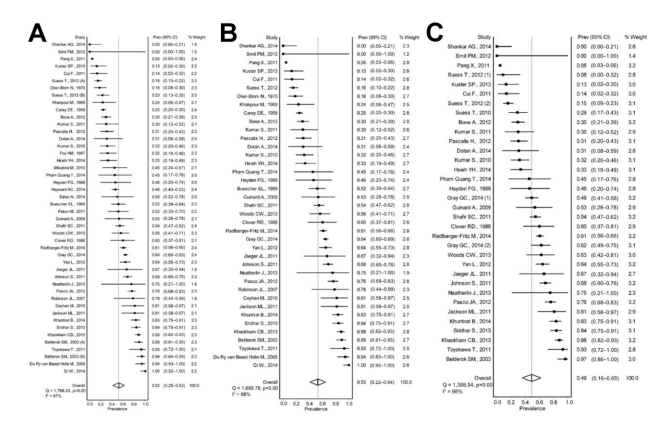


Technical Appendix 2 **Figure 2.** Forest plots for subclinical prevalence of A) influenza overall, B) influenza A, and C) influenza A(H1N1) virus infections generated by using the inverse variance

heterogeneity model for 55 studies included in systematic review and meta-analysis of asymptomatic and subclinical influenza infection prevalence.



Technical Appendix 2 Figure 3. Forest plots for asymptomatic prevalence of A) influenza overall, B) influenza A, and C) influenza A(H1N1) virus infections generated by using the random effects model for 55 studies included in systematic review and meta-analysis of asymptomatic and subclinical influenza infection prevalence.



Technical Appendix 2 Figure 4. Forest plots for subclinical prevalence of A) influenza overall, B) influenza A, and C) influenza A(H1N1) virus infections generated by using the random effects model for 55 studies included in systematic review and meta-analysis of asymptomatic and subclinical influenza infection prevalence.